In vivo human brain imaging at 0.2 T using a whole-body fast field-cycling MRI system

Gareth R. Davies^a, Lionel M. Broche^a, Kerrin J. Pine^b P. James Ross^a and David J. Lurie^a ^a Aberdeen Biomedical Imaging Centre, University of Aberdeen, UK ^b Present address: Imaging Neuroscience, University College London, UK

Purpose: Fast Field-Cycling (FFC) instruments change the main magnetic field strength B_0 during the pulse sequence. With FFC it is possible to obtain image contrast from the dispersion of T_1 over a range of field strengths¹. In a typical pulse sequence the field strength is switched from a polarising field, B_{0p} , to an evolution field B_{0e} , at which relaxation processes of interest occur, before switching to a detection field B_{0d} . FFC requires bespoke magnets, power supplies and ancillary equipment.

Methods: A number of FFC instruments are presented in the literature²⁻⁶. Most are dual magnet designs in which B_{0d} is supplied by one magnet, the second magnet providing offset for B_{0e} . Our magnet (Fig.1) consists of three copper coils, co-wound on a cylindrical former, and potted in epoxy resin (Tesla Engineering Ltd, Storrington, UK). At 2040 mm long, 500 mm bore, it is suitable for human subjects. The magnet has a bare inductance of 5 mH and resistance of 85 m Ω per channel, each requiring 650 A to attain the 0.2 T field specified. The current is supplied by a purpose-built bank of high-power gradient amplifiers (International Electric Co. Oy, Helsinki, Finland).

Results: Fig. 2 shows a transaxial spin-echo FFC image of the brain of a healthy volunteer. Acquistion parameters were: 64x64, field-of-view 300 mm, slice thickness 10 mm, TE 10 ms, TR 1500 ms, field ramp time 20 ms, polarization time 500 ms,

 $B_{0p} = B_{0e} = B_{0d} = 196 \text{ mT} (8.34 \text{ MHz proton frequency}).$



Fig. 1: 0.2 T whole-body FFC imager.



Fig.2: Transaxial image through volunteer's brain.

Discussion and conclusion: Our next step is to employ B_{0e} control to obtain images with T_1 dispersion contrast. We are also working on methods of compensating for environmental magnetic fields, including use of the external correction coils visible in Fig. 1.

References

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